Amendments to the Claims:

Please amend the claims to read as follows:

- 1. (Previously presented) A method for isolating compounds that possess amyloid inhibitory activity from plant matter of the genus *Uncaria*, the method comprising the steps:
 - a) preparing a polar solvent extract of *Uncaria* plant matter, where the polar solvent extraction is selected from one of the extraction methods from the group of extraction methods consisting of extraction with water, extraction with a water solution of a polar alcohol, extraction with a water solution of acetonitrile and extraction with a water solution of another polar organic solvent and running the extract through a first column that comprises hydroxy group containing resin, resin having hydrophobic characteristics but without any hydroxy groups, or a mixture of both;
 - b) eluting the first column with distilled water, followed by eluting with not more than 2-4 column bed volume washings with a dilute polar alcohol/water solution having an alcohol/water ratio not greater than 50/50, and discarding any eluate;
 - c) eluting the first column with one or more column bed volume washings of a polar alcohol/water solution having an alcohol/water ratio between 50/50 and substantially pure alcohol, and collecting and drying the eluted volumes to a dried material.
- 2. (Previously presented) The method of claim 1 wherein the column that comprises hydroxy containing resin, resin having hydrophobic characteristics but without any hydroxy groups, or a mixture of both is a column selected from the group of columns consisting of carbon-containing columns, Tris-acrylate column, LH-20 column, and Affi-prep 10 gel column.
- 3. (Original) The method of claim 1 wherein the polar alcohol/water solution has an alcohol/water ratio of 75/25 or higher.
- 4. (Original) The method of claim 1 wherein the washing in step (c) is effected with substantially pure methanol.
- 5. (Original) The method of claim 1 wherein the plant matter of the genus *Uncaria* is taken from one or more of the plants of the various *Uncaria* species plant group consisting of *tomentosa*,

attenuata, elliptica, guianensis, pteropoda, bernaysli, ferra DC, kawakamii, rhyncophylla, calophylla, gambir, and orientalis.

- 6. (Original) The method of claim 1 wherein the plant matter of the genus *Uncaria* is taken from *Uncaria tomentosa*.
- 7. (Original) The method of claim 6 wherein the *Uncaria tomentosa* plant matter is taken from one or more of the group of plant parts consisting of inner bark and root.
- 8. (Original) The method of claim 1 further comprising the steps:
 - d) applying an aqueous solution of the dried material from step (c) to a second column comprising a hydrophobic resin, the second column having been preparatorily equilibrated in a solvent comprising about 95% water/5% acetonitrile, referred to herein as solvent A, and then eluting the second column with more solvent A and discarding the eluate;
 - e) eluting the second column with a mixture of solvent A containing 10-15% of a solvent comprising about 95% acetonitrile/5% water, referred to herein as solvent B, and collecting and drying the eluted volumes to a dried material.
- 9. (Previously presented) The method of claim 8 wherein the second column comprising a hydrophobic resin is a column selected from the group of columns consisting of C18 SPE, a flash chromatography column, other HPLC columns, and other carbon-containing columns.
- 10. (Currently Amended) The method of claim 1 or 8 further comprising the steps:
 - f) making one or more injections of a solution of the dried material of step (c) or the dried material of step (e) in a solvent selected from the group of solvents consisting of water, water/dilute alcohol and a solvent comprising about 95% water/5% acetonitrile, referred to herein as solvent A, and no more than 10% of a solvent, comprising about 95% acetonitrile/5% water, referred to herein as solvent B, into an HPLC instrument having a diode array uv/vis detector with a graphic display, the HPLC instrument further comprising a reverse-phase column;
 - g) eluting the material through the HPLC column using a solvent gradient profile as follows: 10% solvent B for about the first 20 minutes from start of elution, 10 to 100% solvent B gradient for about minutes 20 to 30 from start of elution, and 100 to 10% solvent

B gradient for about minutes 30 to 32 from start of elution, while observing the uv/vis detector graphic display during the elution gradient over time, and separating fractions of the eluate at elution times corresponding to times associated with the graphic display peaks.

- 11. (Currently amended) The method of claim 10, wherein the reverse-phase column has dimensions of about 2.2cm X 25cm and contains about 95ml of C18 reverse phase resin, wherein the solution of the dried material is a solution of about 50 mg of the dried material of step (c) in about 1-2 ml of solvent A, wherein the step of injecting the solution of dried material into the HPLC may be repeated, wherein a HPLC column solution gradient flow rate is set to about 5 mls per minute, and further wherein the solvent gradient profile is 10% solvent B for 0 to 20 minutes, followed by 10 to 100% solvent B gradient for minutes 20 to 30, and 100% to 10% solvent B gradient from minutes 30 to 31; such that fractions <u>FG</u> though N of the eluate are collected at the following times: fraction G (13-14 minutes), fraction F (15-16 minutes), fraction H (17-20 minutes). fraction I (21 minutes), fraction J (22 - 23 minutes), fraction K1 (24 minutes), fraction K2 (25 minutes), fraction L (26-27 minutes), fraction M (27-28 minutes), and fraction N (28-29 minutes). 12. (Currently amended) The method of claim 10, wherein the reverse-phase column with dimensions of 1.0 cm X 25.0 cm containing 20ml of C18 reverse phase resin, wherein the solution of the dried material of step (c) is a solution of 50 µg of the dried material in 50-100µl of solvent A, wherein the step of injecting the solution into the HPLC is repeated multiple times, wherein a HPLC column solution gradient flow rate is set to about 1.5 mls per minute, and further wherein the solvent gradient profile is 10% solvent B for 0 to 20 minutes, followed by 10 to 100% solvent B gradient for minutes 20 to 30, and 100% to 10% solvent B gradient from minutes 30 to 31; such that fractions FG though O of the eluate are collected at the following times: fraction G (12-13 minutes), fraction F (13-14 minutes), fraction H (15 minutes), fraction I (16 minutes), fraction J (18-19 minutes), fraction K1 (20 minutes), fraction K2 (21 minutes), fraction L (21-23 minutes), fraction M (23 minutes), fraction N (24 minutes), and fraction O (26-27 minutes).
- 13. (Currently amended) The method of claim 10 wherein steps (f) and (g) are as follows:
 - f) injecting a solution of 1 gram of the dried material of step (c) in 5 10 ml of solvent A into an HPLC instrument having a Varian model 320 uv/vis detector set at 230 nm with

- a graphic display, the HPLC further comprising a 4.14 cm X 25 cm Varian Dynamax column further comprising 380 ml of C-18 reverse phase resin, the column fitted to a Varian Prostar 215 solvent delivery system, or the like.
- g) eluting the HPLC column at a solution gradient flow rate of about 50 ml/minute, and further wherein the solvent gradient profile is with a solvent C/solvent D gradient as follows: 0-4 minutes, 25% D; 4-11 minutes, 25-30% D gradient; 11-14 minutes, 30-90% D gradient; 14-17 minutes, 90% D; and 17-19 minutes, 90-25% D gradient, where C is water and D is methanol, such that fractions $F\underline{G}$ through O of the eluate are separated at elution times corresponding to times associated with the graphic display peaks.
- 14. (Currently amended) The method of claim 1 wherein the preparation in step (a) of the extract of *Uncaria* is as follows:
 - 1) adding 4000ml of methanol to 1 kg of *Uncaria tomentosa* and mixing
 - 2) centrifuging the mixture at X2,500g using a centrifuge for 30 minutes and collecting the supernatant;
 - 3) extracting the insoluble material about 3 more times as steps (a) and (b) above;
 - 4) combining the supernatants and evaporating to a dried extract, or to at least about 500 ml volume, using a rotary evaporator at 50°C;
 - 5) washing the dried extract, or the 500ml volume, 4 times with 300ml of petroleum ether, and discarding the ether layer;
 - 6) further evaporating any remaining methanol to dryness using a rotary evaporator at 50°C;
 - 7) extracting the dried extract 5 times with 150ml of distilled water, followed by centrifugation at 2,500Xg for 30 minutes each time, and
 - 8) combining the supernatants and then lyophilizing using a freeze-dryer.
- 15. (Original) The method of claim 14 wherein the further preparation of the extract of *Uncaria* from the resulting lyophilized extract includes the following additional steps:
 - 9) dissolving the resulting lyophilized extract into about 500 ml of distilled water, and applying 50-100ml portions to a 400 ml LH-20 column equilibrated with distilled water.

- 10) eluting the LH-20 column with 1,100ml of distilled water (~3 column volumes) and discarding the amber/yellow, non-active fractions;
- 11) eluting the LH-20 column with 1,100ml of 100% methanol (~3 column volumes) and collecting a set of active fractions and evaporating to dryness using a rotary evaporator at 50°C.
- 16. (Currently amended) The method of claim 8 wherein the aqueous solution of a dried material from step (c) is further prepared by the following steps:
 - 1) dissolving the dried material in water at 80 mg/ml and applying 5 ml at a time to a disposable C18 SPE column (10 gram) equilibrated in a first solvent comprising about 95% water/5% acetonitrile/ 0.1% TFA;
 - 2) washing with 3 column bed volumes of the first solvent and discarding the eluate:
 - 3) eluting with 3 column bed volumes of the first solvent further comprising about 12.5% of a second solvent comprising about 95% acetonitrile/5% water/0.1% TFA; and
 - 4) lyophilizing the corresponding fractions using a freeze-dryer.
- 17. (Currently Amended) The method of claim 8 wherein the aqueous solution of a dried material from step (c) is further prepared by the following steps:
 - 1) dissolving the lyophilized fractions at 5 grams in 20 ml water and applying 20ml at a time to a flash chromatography column;
 - 2) washing with 3 column bed volumes of a first solvent comprising about 95% water/5% acetonitrile/ 0.1% TFA and discarding the eluate;
 - 3) eluting with 3 column bed volumes of the first solvent further comprising about 12.5% of a second solvent comprising about 95% acetonitrile/5% water/0.1% TFA; and
 - 4) collecting and drying the next 3 column bed volumes of eluate.

18-21. (Cancelled)

- 22. (Currently amended) The method of claim 1 further comprising the steps:
 - d) applying an aqueous solution of the dried material from step (c) to a second column, LH-20 or the like, eluting the material from the column with successive column volumes of

water/methanol mixtures containing 0.1% TFA, beginning with 25% methanol and increasing to 100% menthol in 25% increments, and collecting and combining the fractions;

- e) separating, combining and drying a fraction to a dried material, referred to hereafter as compound H, by analytical HPLC, the fraction containing a peak occurring between 7-8 minutes from start of elution on a Dynamax-5μ-C-18 column having dimensions of about 4.6mm X 25cm, using an elution gradient of water for solvent A and methanol for solvent B, A and B each containing about 0.1% TFA, with detection at 280 nm, the gradient conditions being 0 to 9 min fro 25% to 36% B gradient, 3 to 10 min for 36 to100% B gradient, 10 to 12 min for 100 % B and 12 to 13 min for 100 to 25% B gradient, all at a flow rate of about 20 ml/min;
- making one or more injections of a solution of the dried material of step (e) above in a solvent comprising water/methanol 80/20 containing about 0.1% TFA and applied at about 150 mg/run to a preparative HPLC Dynamax 5μ-C-18 column with dimensions of about 21.4mm X 25cm, using substantially the same elution gradient as used in step (e) above, with detection at 280 and 300 nm, the gradient conditions being 0 to 3 min for 20% to 25% B gradient, 3 to 9 min for 25 to 45% B gradient, 9 to 10 min for 45 to 100% B gradient, 10 to 12 min for 100% B and 12 to 13 min for 100 to 25%B gradient, all at a flow rate of about 20 ml/min, the a compound H fraction eluting between 7-8 minutes from start of elution, and;
- g) repeating steps (e) and (f) above until the peak as seen on analytical HPLC in step (e) is relatively pure.

23-38. (Cancelled)

- 39. (Currently amended) The method of claim 1 wherein the polar organic solvent of step (a) is selected from the group of polar organic solvents consisting of triethanolamine; and acetone.
- 40. (Previously presented) The method of claim 2 wherein the carbon-containing columns are selected from the group consisting of C2 column, C4 column and C18 column.